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Principal Investigator Dr. Louis A. Jones

Department of Chemistry

North Carolina State University

Raleigh, North Carolina 27607

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Acenaphthene III. Coupling Reactions. The Preparation of 4,4'-Dinaphthalic Anhydride and 4',1-Naphthylnaphthalic Anhydride¹

LOUIS A. JONES² AND RONALD WATSON

Department of Chemistry, North Carolina State University, Raleigh, N.C. 27607 Received October 31, 1972

The syntheses of the previously unknown 4,4'-dinaphthalic anhydride (4) and 4',1-naphthylnaphthalic anhydride (5) is reported. The vinylic bromination of 5-(1,-3,4-dihydronaphthyl)acenaphthene (11) is also described.

On décrit la synthèse des anhydrides dinaphtalique-4,4'(4) et (naphtyl-1)-4 naphtalique (5) inconnus jusqu'à maintenant. On rapporte aussi les résultats obtenus lors de la bromination vinylique du (dihydro-3,4 naphtyl-1)-5 acénaphtène (11). [Traduit par le journal]

Can. J. Chem., 51, 1833 (1973)

Recent investigations (1, 2) have resulted in the preparation of 4,5-diaminonaphthalic anhydride (1) and 3,4-diaminonaphthalic anhydride (2). As part of a continuing study in the chemistry of acenaphthene (3) and its derivatives, we now report the syntheses of 4,4'-dinaphthalic anhydride (4) and 4',1-naphthylnaphthalic anhydride (5).

Thallium bromide (1) has been used as a reagent for the syntheses of biaryls from aromatic Grignard reagents (3). Consequently, 5-acenaphthyl magnesium bromide, prepared from 5-bromoacenaphthene (4), was treated with thallium bromide to give 5,5'-diacenaphthene (6) in low yield. Alternatively it was found that the

same product could be obtained in far better yield by utilizing the coupling technique of Kharasch and Fields (5) in which cobalt chloride is used as a catalyst. The fact that coupling has indeed taken place in the *peri*-position was established from consideration of its n.m.r. spectrum. The bridgehead protons gave a sharp singlet at τ 6.57 as did the aromatic protons 6,6′, 7,7′, and 8,8′ at τ 2.74. The remaining protons exhibited an AB quartet with a J value of 7.3 Hz. Oxidation to the dianhydride 4 was achieved by treatment of the product with sodium dichromate in glacial acetic acid.

Garvey and his co-workers (6) have previously reported the synthesis of 5,5'-diacenaphthene by a different procedure, but no experimental details or spectral data were given.

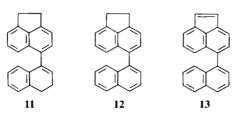
When the dimer 6 was treated with 2,3-dichloro-5,6-dicyano-1,4-benzoquinone (DDQ) dehydrogenation took place to give 5,5'-diacenaphthylene (7) which readily consumed 2 mol of bromine to give the tetrabromo-compound 8. Also, oxidation with sodium dichromate again gave the dianhydride 4.

When the Grignard reagent from 5-bromoacenaphthene was treated with α -tetralone in tetrahydrofuran, the expected coupled product 9was only obtained in low yield. Rather, we obtained acenaphthene and the aldol condensation product 10 of α -tetralone which was identified by its elemental analysis and by preparation of its 2,4-dinitrophenylhydrazone. This same product has previously been isolated (7) from the reaction of α -tetralone with cyclohexyl magnesium bromide. Consequently, treatment of α tetralone with 5-lithio-acenaphthene (from 5-

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²Author to whom correspondence should be addressed.

bromo acenaphthene and *n*-butyl-lithium) resulted in the isolation of the hydroxy compound **9** in far better yield. Dehydration to 5-(1,-3,4-dihydronaphthyl)acenaphthene (**11**) was readily effected by treatment with polyphosphoric acid. This product was subsequently dehydrogenated by 10% palladium-on-charcoal to 5',1-naphthylacenaphthene (**12**) or by DDQ which gave 5',1-naphthylacenaphthylene acenaphthylene (**13**). Treatment of either compound **12** or **13** with sodium dichromate in glacial acetic acid resulted in oxidation to 4',1-naphthylnaphthalic anhydride (**5**).



Nitration of this anhydride resulted in the isolation of a yellow solid. Consideration of its elemental analysis and mass spectrum indicated the presence of two nitro-groups. Although the positions of these nitro-groups have not as yet been unambiguously assigned, previous experience (8) in this laboratory has shown that, in the case of nitration of 1,1'-binaphthyl the *peri*positions are more susceptible to nitration. Consequently we propose that this dinitro-compound has the structure 14. Further, its n.m.r. spectrum exhibits two aromatic multiplets in the ratio of 4:6 which is the pattern one would expect for this structure.

The possibility of introducing a bromine atom in the 3'-position of the system 11 and then dehydrobromination as an alternate route to naphthyl acenaphthene (12) led us to attempt allylic bromination utilizing N-bromosuccinimide (NBS) in carbon tetrachloride as the brominating media (9). This resulted in an inseparable mixture of isomers. When dimethylformamide (DMF) was used as solvent, the reaction proceeded smoothly to give a good yield of a mono-brominated product. This product, however, did not appear to be the required allylically brominated system 15 since it gave no precipitate with alcoholic silver nitrate at room temperature. Consideration of its n.m.r. spectrum revealed that bromination had in fact taken place in the vinyl-2'-position to give compound 16, the main feature of the spectrum being the disappearance of the triplet at τ 4.0 in the starting material and the appearance of a singlet for the 3' and 4'-protons. The structure has been unambiguously confirmed by a single crystal X-ray analysis (10).

Although rare, instances of vinylic bromination in NBS reactions are not altogether unique (11, 12). Pines and his co-workers (13) found that bromination of α -methylstyrene with NBS in carbon tetrachloride gave, as well as the expected 2-phenylallyl bromide (17), 1-methyl-1-phenylvinyl bromide (18), in a ratio of 3:1. In an earlier study, Roberts and Trumbell (14) found that the reaction of camphene with NBS in carbon tetrachloride gave a 35% yield of mono-brominated

$$C=CH_2$$
 $C=CHB_1$ CH_2B_1 CH_3 CH_3

products, of which 65% proved to be the vinylic substituted product 8-bromocamphene. In each case a free radical mechanism is proposed.

In our hands, the reaction of α-methylstyrene with NBS in DMF gave 56% of mono-brominated products. From g.l.c. studies of the crude

mixture and the pure isomers it was shown that the mixture consisted of 80% of 17 and 29% of 18. The same proportions were obtained from consideration of the n.m.r. spectra of both the crude mixture and the pure compounds. It had previously been reported (9b) that NBS was inactive towards styrene and that no monobrominated products were obtained. However, reaction of styrene with NBS in DMF as the solvent was found to produce a 30% yield of dibromostyrene. This would seem to indicate, therefore, that in DMF the addition of bromine to the double bond takes preference over substitution, although in the case of α -methylstyrene, no dibromo-substituted products were detected. This was confirmed by the fact that when the reaction of compound 11 with NBS in DMF was carried out in the dark, the same product, 16, was obtained in identical yield as had been obtained previously. Further, addition of bromine to the dihydro system 11 again gave the vinylic brominated product 16, also in the same yield. The mechanism for the formation of 16 therefore, evidently involves an initial addition of bromine to the double bond to form a dibromointermediate, which then spontaneously dehydrobrominates to give the product 16.

This is consistent with the findings of Horner and Winkelmann (9d) that with increasing polarity of the solvent used in the bromination reaction of NBS the allylic bromination activity disappears and the addition of bromine to the double bond becomes the main reaction.

Experimental³

5,5'-Diacenaphthene (6)

5-Bromoacenaphthene (10.0 g) in anhydrous tetrahydrofuran (100 ml) was added to magnesium turnings (1.98 g) in tetrahydrofuran (100 ml) and the mixture heated to reflux under nitrogen for 16 h. The resulting solution of 5-acenaphthyl magnesium bromide was added to 5-bromoacenaphthene (10.08 g) and anhydrous cobalt chloride (9.5 g) in anhydrous tetrahydrofuran (100 ml) and the mixture heated to reflux for a further 16 h. The mixture was then poured on ice, acidified (diluted HCl), and extracted with methylene chloride. The organic layer was separated, washed, dried, and evaporated to leave a

brown semi-solid which was chromatographed on alumina. Elution with hexane gave unreacted bromoacenaphthene and some acenaphthene followed by the product 6 (3.2 g, 49%) which was recrystallized from methylene chloride – hexane to give pale yellow needles, m.p. 176–178°, $v_{c=c}$ 1610 cm⁻¹ (Nujol); τ (CDCl₃) 2.46–2.72 (4H, AB quartet, J=7.3 Hz, Ar), 2.74 (6H, s, Ar), and 6.57 (8H, s, CH₂).

Anal. Calcd. for C₂₄H₁₈: C, 94.1; H, 5.8. Found: C, 94.2; H, 5.9.

4,4-Dinaphthalic Anhydride (4)

Diacenaphthene (3.3 g), sodium dichromate (27 g), and glacial acetic acid (500 ml) were refluxed overnight, cooled, and added to water (600 ml). The precipitated solid was collected and recrystallized from dimethylformamide to give 4 (3.2 g, 71%) as a yellow amorphous solid m.p. 362° (dec.), v_{max} (Nujol) 1780, 1730 (C=O) and 1595 cm⁻¹ (C=C).

Anal. Calcd. for $C_{24}H_{10}O_6$: C, 73.10; H, 3.45%. Found: C, 72.85; H, 3.15.

5,5'-Diacenaphthylene (7)

Diacenaphthene (1.73 g), DDQ (3.12 g), and benzene (40 ml) were heated at reflux overnight; the mixture was cooled, diluted with hexane (200 ml) and the hydroquinone, which had formed during the reaction, was removed by filtration. The filtrate was chromatographed on alumina. Hexane eluted 7 (1.17 g, 70%), yellow plates, m.p. 164–165% (from ethanol), τ (CDCl₃) 2.34–2.76 (10H, m, Ar), and 3.02 (4H, s, methine).

Anal. Calcd. for $C_{24}H_{14}$: C, 95.35; H, 4.65. Found: C, 95.25; H, 4.5.

1,1',2,2'-Tetrabromo-5,5'-diacenaphthene (8)

To diacenaphthylene (1.5 g) in hexane (130 ml), a solution of bromine (1.6 g) in hexane (20 ml) was added dropwise and the mixture stirred for 8 h at 20%. The mixture was concentrated and the precipitated solid separated by filtration and recrystallized to give 8 (1.64 g, 50%), as a yellow amorphous solid, m.p. 218–219° (dec.) (from benzene).

Anal. Calcd. for C₂₄H₁₄Br₄: C, 46.3; H, 2.25; Br, 51.45. Found: C, 46.45; H, 2.10; Br, 51.35.

Reaction of 5-Acenaphthyl Magnesium Bromide with α-Tetralone

A solution of α-tetralone (6 g) in anhydrous tetrahydrofuran (50 ml) was added to the Grignard solution obtained from 5-bromoacenaphthene (10 g), magnesium turnings (1.08 g), and anhydrous tetrahydrofuran (100 ml) and the mixture refluxed under nitrogen for 3 days; it was then poured onto ice, acidified (dilute HCl), and extracted with methylene chloride. The extract was washed (NaHCO₃ then water) and dried (Na₂SO₄), to give a residue which was chromatographed on Florisil. Hexane eluted acenaphthene (3 g, 45%) and further elution with ether gave a brown gum. Trituration of this gum with ether gave 10 (1.9 g, 32%), pale yellow crystals, m.p. 132-133° (from methylene chloride - hexane) (lit. (7) m.p. 132-133°). Evaporation of the filtrate gave the required product 9 (2.5 g, 19%) as a pale brown amorphous solid. m.p. 112-114° (from methylene chloride - hexane), v_{max} (Nujol) 3450 (OH), 1612 cm⁻¹ (C=C).

Anal. Calcd. for C, 88.0; H, 6.65. Found: C, 88.25; H, 6.5.

³Melting points were determined on a Thomas-Hoover capillary melting point apparatus and elemental analyses performed by Galbraith Laboratories, Inc., Knoxville, Tennessee. I.r. spectra were measured on a Beckman I.R.-33 i.r. spectrophotometer and n.m.r. spectra were obtained from a Varian HA-100 spectrometer.

1-Hydroxy-1,2,3,4-tetrahydro-5',1-naphthylacenaphthene (9)

5-Bromoacenaphthene (50 g) in ether (300 ml) and a hexane solution of n-butyl-lithium were mixed at 20° and stirred at this temperature under nitrogen for 20 min. α -Tetralone (31 g) in ether (150 ml) was then added and the mixture was stirred for a further 15 h at 20° . The mixture was then poured onto ice and acidified (dilute HCl). The ether layer was separated off, washed, dried, and evaporated to leave a light brown semi-solid. This was triturated with hexane and the residue crystallized from methylene chloride – hexane. The product 9 (40 g 63%) formed a fawn-colored semi-crystalline solid, m.p. $112-114^{\circ}$.

5-(1,-3,4-Dihydronaphthyl) acenaphthene (11)

The preceding product (30 g) was added with stirring to polyphosphoric acid (500 g) and the mixture was stirred at 60° for 5 h; it was then poured into ice-cold water (1000 ml) and extracted with ether. The extract was washed with aqueous sodium bicarbonate solution and then with water, dried (Na₂SO₄), and evaporated. The dehydrated product 11 (27 g, 95%) formed light brown crystals, m.p. 112° (from methylene chloride – hexane), v_{C=C} (Nujol) 1595 and 1610 cm⁻¹; τ (CDCl₃) 2.64–3.50 (9H, m, Ar), 4.0 (1H, t, 2'-H), 6.76 (4H, s, 1-H and 2-H), 7.04–7.32 (2H, m, CH₂), and 7.5–7.8 (2H, m, CH₂).

Anal. Calcd. for C₂₂H₁₈: C, 93.6; H, 6.4. Found: C, 93.4; H, 6.2.

5',1-Naphthylacenaphthene (12)

The dihydro-compound 11 (16.5 g) and 10% palladium-on-charcoal (5 g) in α -methylnaphthalene (250 ml) were heated to reflux under nitrogen for 25 h; the mixture was cooled, filtered, the α -methylnaphthalene distilled off under reduced pressure, and the residue chromatographed on alumina. Hexane-benzene (1:1) eluted the product (14.5 g, 89%) as pale yellow crystals, m.p. 103–104° (from hexane), τ (CDCl₃) 2.02–2.84 (12H, m, Ar), and 6.59 (4H, s, 1-H and 2-H).

Anal. Calcd. for $C_{22}H_{16}$: C, 94.3; H, 5.7. Found: C, 94.1; H, 5.8.

5', I-Naphthylacenaphthylene (13)

The dihydro-compound 11 (22 g), DDQ (37 g), and benzene (300 ml) were heated to reflux overnight and then worked-up as for diacenaphthylene. The dehydrogenated product (12 g, 55%) afforded yellow crystals, m.p. $112-114^{\circ}$ (from ethanol), τ (CDCl₃) 2.1-2.9 (12H, m, Ar), and 3.0 (2H, s, 1-H and 2-H).

Anal. Calcd. for C₂₂H₁₄: C, 94.95; H, 5.05. Found: C, 94.7; H, 5.1.

4',1-Naphthylnaphthalic Anhydride (5)

The preceding product (11.7 g), sodium dichromate (40 g) and glacial acetic acid (500 ml) were heated to reflux overnight, cooled, and added to water (1000 ml). The precipitated solid was collected and recrystallized to give 5 (10 g, 74%) as dark yellow crystals, m.p. 250-252° (from aqueous acetic acid), $v_{\rm CO}$ (Nujol) 1770, 1730 cm⁻¹; τ (CDCl₃) 1.24-1.47 (2H, m, Ar), and 1.95-2.84 (10H, m, Ar).

Anal. Calcd. for $C_{22}H_{12}O_3$: C, 81.5; H, 3.75. Found: C, 81.25; H, 3.75.

Nitration of 4',1-Naphthylnaphthalic Anhydride

To a stirred solution of compound 5 (1.4 g) in methylene dichloride (150 ml), 90% nitric acid (15 ml) was added dropwise and the mixture heated to reflux for 60 h; it was then poured into water (300 ml), the organic layer separated, washed (NaHCO₃, then water), dried (MgSO₄), and evaporated to give a pale yellow solid. This was dissolved in methylene dichloride and addition of hexane gave the product 14 (0.6 g, 32%) as an amorphous yellow solid, (m/e 413) m.p. 280° (dec.) (from acetic acid), v_{max} (Nujol) 1775, 1730 $v_{c=0}$, 1530, and 1355 cm⁻¹ (NO₂); τ ((CD₃)₂SO) 1.15–1.56 (4H, m, Ar), and 1.69–2.26 (6H, m, Ar).

Anal. Calcd. for $C_{22}H_{10}N_2O_7$: C, 63.75, H, 2.4; N, 6.75. Found: C, 63.75; H, 2.3; N, 6.65.

2-Bromo-3,4-dihydro-5',1-naphthylacenaphthene (16)

(a) To compound 11 (3 g) in dimethylformamide (100 ml) a solution of NBS (1.89 g) also in dimethylformamide (50 ml) was added and the mixture stirred overnight at 20°; it was then poured into water (700 ml) and extracted with ether. The extract was washed with water, dried (MgSO₄), and evaporated to give a pale yellow gum which was chromatographed on Florisil. Hexane eluted the product 16 (2 g, 55%), which gave yellow crystals, m.p. 150–152° (from methylene chloride – hexane), $v_{C=C}$ (Nujol) 1604, 1585 cm⁻¹; τ (CDCl₃) 2.68–3.66 (9H, m, Ar), 6.74 (4H, s, 1-H and 2-H), and 7.07 (4H, s, 3'-H and 4'-H).

Anal. Calcd. for $C_{22}H_{17}Br: C$, 73.15; H, 4.7; Br, 22.15. Found: C, 72.95; H, 4.5; Br, 22.3.

(b) To compound 11 (6 g) in carbon tetrachloride (100 ml) a solution of bromine (3.5 g) in carbon tetrachloride (20 ml) was added dropwise and the mixture stirred at room temperature for 30 min; evaporation of the solvent gave 16 (4.3 g, 56%) as light brown crystals, m.p. 150-152° (from methylene chloride – hexane).

Styrene with N-Bromo Succinimide

A solution of NBS (50 g) in dimethylformamide (80 ml) was added to styrene (100 ml) in dimethylformamide (20 ml). After the initial exothermic reaction had subsided, the mixture was stirred overnight at 20°. The solvent and excess styrene were removed under reduced pressure and the residue chromatographed on silicic acid. Hexane eluted dibromostyrene (21.3 g, 30%) as white needles m.p. 70-72° (lit. (15) 72-73°) (from aqueous ethanol).

a-Methylstyrene with N-Bromosuccinimide

A solution of NBS (100 g) in dimethylformamide (250 ml) was added to α-methylstyrene (200 ml). After the initial exothermic reaction had subsided the mixture was stirred overnight at room temperature, poured into water, and extracted with methylene chloride. The extract was washed with water, dried (MgSO₄), and evaporated to give a dark brown liquid. Excess α-methylstyrene was removed under reduced pressure and vacuum distillation of the residue gave the mono-brominated products (61.6 g, 56%), b.p. 95–105°/11 mm, a yellow liquid containing a mixture of isomers. The mixture (13 g) was chromatographed on silicic acid and eluted with hexane. The first fractions were composed of pure 1-methyl-1-phenyl-vinyl bromide (18) while the end fractions were composed

of pure 2-phenylallyl bromide (17) (11). From g.l.c. studies of the pure isomers it was calculated that the crude mixture of bromides was composed of 80% of 17, τ (CDCl₃) 2.4–2.76 (5H, m, Ar), 4.49 (1H, s, vinyl-H), 4.56 (1H, s, vinyl-H) and 5.68 (2H, s, CH₂); and 20% of 18 τ (CDCl₃) 2.69 (5H, s, Ar), 3.57 (1H, m, vinyl-H), and 7.83 (3H, s, CH₃).

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Freparation of 4,4',5,5',-Tetranitro-1,1'-binaphthyl

Louis A. Jones¹ and Charles T. Joyner

Department of Chemistry, North Carolina State University, Raleigh, N.C. 27607

The synthesis and proof of structure of 4,4',5.5'-tetranitro-1,1'-binaphthyl is described.

Previous communications from this laboratory have described the nitration products of acenaphthene and the subsequent conversion to diaminonaphthalic anhydrides or derivatives thereof (10, 11). As an extension of these studies, the nitration of binaphthyl (Compound 1) was examined since it differed from acenaphthene in that the 8-position of the naphthalene moiety was available for substitution, whereas the corresponding position in acenaphthene was not. Lossen reported in 1867 that the direct nitration of compound 1 with furning nitric acid resulted in a product whose empirical formula was C20H10-(NO2)4 although no structural assignment was reported (15). Mild nitration of compound 1 has been reported to yield the 4,4'-dinitro derivative (Compound 2) (19). In view of the paucity of information concerning this nitration, we became interested in determining the products obtained under a variety of nitrating conditions and, in particular, attempting to prepare 4,4',5,5'-tetranitro-1,1'binaphtnyi (Compound 3), since this could be converted to a tetraamino compound capable of forming ladder polymers with the appropriate acids or anhydrides (3). Although unsymmetrical tetranitrobinaphthyl compounds have been prepared by the Ullman reaction with dinitrohalonaphthalenes (17), Compound 3 has not been previously reported.

To prepare the necessary dinitronaphthalene derivative for Ullman coupling, 1-bromo-5-nitronaphthalene (Compound 4) was nitrated under vigorous conditions and yielded two products. Liquid chromatographic separation and analysis showed the mixture to be 80% of the desired 1-bromo-4,5-dinitronaphthalene (Compound 5) and surprisingly, 1,5-dinitronaphthalene (Compound 6). Only one other unusual displacement of this type has been reported (12). The nmr of compound 5 indicated that the shielding effect of the bromine and the nitro group are essentially equal since protons 2 and 3 had the same chemical shifts and appeared as a single 2H peak at δ 8.2. Alternatively, nitration of 1-bromo-4-nitronaphthalene (Compound 7) produced the desired Compound 5 in high yields (8) with no noticeable replacement side products and the ir and nmr of Compound 5 produced by the two methods were identical.

Rogavik et al. reported that when 1,8-dinitronaphthalene was heated with phthalic anhydride in acetic acid and iron, a phthaloperinone was formed (18). Examination of Dreiding models showed the formation of such a compound could occur only when the nitro groups are in the 3,4- or 4,5-positions. However, the directive properties of the hitro group and the naphthalene moiety suggest that 3,4-substitution would be most unlikely (7, 10, 16). Thus when Compound 5 from both sources was treated with phthalic anhydride and iron in acetic acid, the 3-bromo [12H-benz[4,5]isoindolo[2,1a]perimidin-12-one] (Cempound 8) was formed in good yield, offering further support for the 4,5-dinitro derivative 5.

When Compound 5 was heated with activated copper (21) to 180°C under nitrogen, complete decomposition

Figure 1. Preparation of 4,4',5,5'-tetranitro-1,1'-binaphthyl and derivatives

occurred while the use of refluxing solvents such as nitrobenzene, ethylene glycol, and butyl cellosolve for long periods of time resulted in the recovery of unreacted Compound 5. The literature contains a few examples of the use of highly polar aprotic solvents to obtain fair yields of coupled benzene products via the Ullman reaction (1. 13). However, when Compound 5 was refluxed for 24 hr with activated copper (21) in distilled N,N-dimethylformamide (DMF), filtered, and the mixture then quenched with water, a red precipitate was obtained. Elemental analysis indicated an empirical formula of C12H11N3O4 while the mass spectrum showed a molecular ion of 261. The nmr contained a singlet in the aliphatic region at δ 3.0 which integrated to six protons, and the data are consistent with 1-N, N-dimethylamino-4,5-dinitronaphthalene (Compound 9) which was obtained in 89% yield. It is interesting to note, however, that when 1-bromonaphthalene is refluxed with Cu(1) in DMF, it is not the N,Ndimethylamino derivative that is produced but rather the reduced product naphthalene (2). It is apparent that two different mechanisms are operating but insufficient data are available to postulate a mechanism for either reaction. The use of other solvents such as dimethylsulfoxide, dimethylacetamide, and N-methylpyrrolidin-2-one, resulted in the production of 1,8-dinitronaphthalene (Compound 10) suggesting that acidic α -hydrogens are required for this reduction although the reaction in DMF is anomalous. Similar replacement results were obtained

¹ To whom carrespondence should be addressed.

with compound 7 was used with the above corresponding solvents.

Several alternative synthetic routes were then studied to a fect the preparation of Compound 3. Compound 1 was propared by either Uriman-type coupling (6, 17) of 1-lead numbratione (Compound 11) or the Busch reaction ,5) of 1-bremenaphtralene (Compound 12). The reaction of Compound 1 with a mixture of acetic acid-nitric acid gave a compound which was identical to that prepared in fair yield from 1-rodo-4-nitronaphthalene (Compound 13) and eagger powder under standard Ullman conditions (Figure 1). When Compound 1 was treated with a mixture of nitric acid and sulfuric acid at room temperature and then refluxed, there was obtained a 53% yield of a compound whose elemental and mass spectral analysis (M/e)= 434) corresponded to $C_{20}H_{10}(NO_2)_4$ or Compound 3. The nmr spectrum of Compound 3 exhibited two multipiets at δ 7.6-8.1 and δ 8.3-8.7 which integrated to 6:4 protons. The area of the latter multiplet can be attributed to the 3,3',6,6' protons while the former arises from the 2.2',7.7',8,8' protons. Similar yields of the same product were obtained using acetyl nitrate (4) or nitronium tetrafluoroporate (14) as the nitrating agents on Compound 2, but direct nitration of Compound 1 was superior method.

Although it was assumed on the basis of physical evidence that the product thus obtained was the desired Compound 3, chemical support was sought for confirmation. Thus, when Compound 3 was treated with phthalic anhydride and iron in acetic acid (vide supra) the analogous bisphthaloper.none binaphthyl compound, 3,3'bi[12-benz(4,5)-isoindolo(2,1-a)perimidin-12-one] was obtained. One of the three possible geometric isomers is shown.

Compound 14

Elemental analysis supported the empirical formula of $G_{38}H_{18}N_2O_2$ and the mass spectrum showed a molecular fon of 533. The nmr contained a complex aromatic proton pattern and no nitro group absorption was present in the ir spectrum although the carbonyl group stretching frequency (1690 cm⁻¹) was observed. Further, when Compound 3 was reduced and the resulting tetraamine (Compound 15) reacted with pathalic analysis of Compound 14 was produced. The spectral analysis of Compound 15 indicated a molecular ion of 314 and N-H absorption in both the nmr and ir spectra, further supporting assigned structure of Compound 14.

Experimental

Meiting points were determined on a Thomas Hoover Capillary meiting point apparatus, and elemental analyses were performed on a Perkin-Elmer Model 240 analyzer or by Galbra'th Laboratory, Inc., Knoxville, Tenn. The nmr spectra were run on a Varian HA 100 NMR Spectrometer with followed pulsariane as an internal standard, and in spectra were obtained from either a Perkin-Elmer 521 or 25% in outcometer. An INEL Model MG12 produced the man outcometer. An INEL Model MG12 produced the

This is taking were obtained from commercial sources: 3.1'-bin, pritry: $\sqrt{\text{Compound}}$ 1) 1-econopythalene (Com-

pound 11), 1-bromonaphthalene (Compound 12), nitronium tetrafluoroborate, palladium on calcium carbonate, and palladium black. Bakerflex silica gel 1B was used for the tlc work, and deuterated solvents were supplied by Stohler isotopes.

The activated copper powder was prepared as described by Vogel (21), 4,4'-Dinitro-1,1'-binaphthyl was synthesized from 1,1-binaphthyl and 1-iodo-4-nitronaphthalene (19) and 1,5-dinitrophthalene was prepared according to the method of Friedlander (8).

4.4',5.5'-Tetranitro-1,1'-binaphthyl (Compound 3). Method A. To a slurry of Compound 1 (10.0 grams) in glacial acetic acid (50 ml) and sulfuric acid (50 ml) were added 30 ml of 90% nitric acid (sp gr 1.5) in small portions with vigorous stirring keeping the temperature below 30°C. When addition was complete, the reaction was refluxed for 24 hr and then cooled to 0°C. The precipitate was collected, and two recrystallizations from glacial acetic acid gave 9.0 grams (53%) of Compound 3 as light cream-colored needles, mp 320–330°C dec; ir (nujol) 1535 cm⁻¹ (NO₂); mass spectrum m/e 434 (molecular ion); nmr (DMSO- d_6) δ 7.6–8.1 (6H multiplet, aromatic), and δ 8.3–8.7 (4H multiplet, aromatic).

Anal. Calcd $C_{20}H_{10}N_4O_8$: C, 55.31; H, 2.32; N, 12.90. Found: C, 55.17; H, 2.32; N, 12.74.

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Method B. Two grams of Compound 2 were refluxed for 28 hr in a solution of acetic acid (50 ml), sulfuric acid (10 ml), and 90% nitric acid (25 ml). The reaction mixture was cooled and 1.8 grams (50%) of Compound 3 were collected and recrystallized from acetic acid. The ir and nmr spectra of this compound were identical to those of the material prepared by direct nitration of Compound 1.

Method C. Five grams of Compound 1 were added with rapid stirring to NO_2BF_4 (5.0 grams) in freshly distilled tetrahydrothiophene-1,1-dioxide (50 ml). The reaction mixture was heated to $80^{\circ}C$ for 1 hr, poured into water (50 ml), the precipitate collected and recrystallized from acetic acid to yield 3.9 grams (45%) of product, which had the same ir and nmr spectra as that prepared by Method A.

Method D. A suspension of Compound 1 (2.0 grams) in acetic anhydride (10 ml) was added slowly to a solution of acetyl nitrate (4) in acetic anhydride at -20° C. After 10 min of stirring, the temperature was raised to 20° C and maintained for 10 min. Finally, the temperature was raised to 80° C for 2 hr during which time dark brown fumes were evolved. The reaction mixture was cooled, added to water (200 ml), and the precipitate filtered and recrystallized from acetic acid to yield 1.8 grams (50%) of Compound 3. Ir and nmr spectra showed this compound to be identical to that prepared by Method A.

1-Bromo-5-nitronaphthalene (Compound 4). Over a 5-hr period, bromine (85 grams) was added to a rapidly stirred melt of 1-nitronaphthalene (20) (173 grams) at 80° C. When all of the bromine had been added, the temperature was raised to 100° C and stirring was continued for 2 hr to expel excess bromine and HBr. The reaction product was recrystallized from ethanol to yield 213 grams (85%) of Compound 4 as orange-yellow needles: mp 122.0-123.0°C [lit. (20) mp 123°C] ir (nujcl 1535, 1350 cm⁻¹ (NO₂); nmr (CDCl₃) δ 7.3-8.5 (multiplet, aromatic).

1-Bromo-4,5-dinitronaphthalene (Compound 5). Method A. To a sturry of Compound 4 (252 grams) in 1500 mt of acetic acid and 150 mt of sulfuric acid, 150 mt of nitric acid (90%) were added dropwise with rapid stirring. Then the reaction mixture was heated to 80°C for 30 min; 250

ml of nitric acid (90%) were added, and the mixture was refluxed for 22 hr. Cooling slowly to 20°C precipitated the product as very pale yellow-white needles. A second crop was precipitated by adding the filtrate to water (3 liters). Both crops were combined and recrystallized from acetic acid to yield 240 grams of cream-colored needles; mp 156-158°C. Qualitative tic developed in ether-hexane (2:1 v/v) showed two compounds present.

One gram of the mixture was chromatographed on Florisil (250 grams) and eluted with ether-hexane (3:1 v/v). The first fraction was identified as Compound 6 by mixed melting point and comparison of its ir and nmr spectra with those of an authentic sample. A total of 0.19 grams was recovered: mp 218°C. The second fraction yielded a total of 0.78 grams of Compound 5. The reaction yield based on the amount of Compound 4 produced was 65%; mp 176.0°C [lit. (20) mp 174°C]; ir (nujol) 1535, 1350 cm⁻¹ (NO₂); nmr (CDCl₃) δ 7.8-8.0 (1H triplet, aromatic), δ 8.5-8.7 (1H doublet, aromatic), δ 8.2 (2H singlet, aromatic), and δ 8.3-8.4 (1H doublet, aro-

1,5-Dinitronaphthalene (Compound 6). Twenty-five ml of 90% nitric acid were slowly added to a suspension of 1-nitronaphthalene (20) (20.0 grams) in acetic acid (100 ml) and sulfuric acid (50 ml). After the addition, the mixture was refluxed for 5 hr and then cooled to 10°C. The precipitated product was recrystallized from acid to yield 23.0 grams (91%) of Compound 6 as pale yellow-white needles: mp 219.5°C [lit. (8) mp 219°C]; ir (nujol) 1350. 1520 cm⁻¹ (NO₂); nmr (CDCl₃) δ 7.6-7.8 (2H triplet, aromatic), δ 8.1-8.3 (2H doublet, aromatic) and δ 8.6-8.8 (2 H doublet, aromatic).

1-Bromo-4-nitronaphthalene (Compound 7). To a solution of Compound 12 (20.7 grams) in 70 ml of acetic acid was added 50 ml of nitric acid (sp gr 1.42) at a moderate rate with rapid stirring. After refluxing for 2 hr the reaction solution was cooled to 0°C and the first crop of crystals collected. The filtrate was then added to 500 ml of water and a second crop of precipitate was recovered. Both crops were recrystallized from ethanol to yield a total of 20.5 grams (81.4%) of Compound 7 as light yellow needles: mp 86.5-87.0°C [lit. (9) mp 87°C]; ir (nujol) 1530, 1345 cm $^{-1}$ (NO₂); nmr (CDCl₃) δ 7.6–8.6 (multiplet, aromatic).

3-Bromo[12H - Senz[4,5]isoindolo[2,1-a]perimidin-12one] (Compound 8). Phthalic anhydride (1.5 grams), powdered iron (3.4 grams), Compound 5 (3.0 grams), and acetic acid (50 ml) were refluxed for 4 hr and the reaction mixture filtered while hot. As the filtrate cooled, a red-brown precipitate formed, which was collected and recrystallized from acetic acid to yield 2.9 grams (75%) of Compound 8 as maroon needles: mp 243.5-245.5°C; ir (nujol) 1690 cm $^{-1}$ C=-O); nmr (DMSO-d₆) δ 7.1-8.1 (complex of multiplets, aromatic).

Anal. Calcd C₁₆H₉N₂OBr: C, 61.89; H, 2.60; N, 8.02. Found: C, 61.75; H, 2.61; N, 7.91.

1-N,N-Dimethylamino-4,5-dinitronaphthalene pound 9). Ten grams of Compound 5 and activated copper (10 grams) were reacted in refluxing distilled DMF (30 ml) for 24 hr. The reaction mixture was then poured into 100 ml of cold water and the precipitate collected and extracted (Soxnlet) with methanol. The extract was reduced to 50 ml and cooled to 0°C to precipitate 9.0 grams (89%) of the product as red-brown needles: mp 775.0-175.5°C [lit. (20) mp 176°C]; ir (nujol) 1560 cm=1 (C-N); mass spectrum m/e 261 (molecular ion); nmr $(CDCl_5)$ δ 3.6 (EH singlet, methyl), δ 7.0-7.7 (2H multiplet, aromatic) 58.1-8.5 (31) multiplet, aromatic).

Anal. Calcd C₁₂H₁₁N₃O₄; C, 55.17; H, 4.24; N, 16.08. Found: C, 55.07; H, 4.26; N, 16.12.

1,8-Dinitronaphthalene (Compound 10). Ten grams of (Compound 5) and activated copper (10 grams) were reacted in DMAC (50 ml) at reflux for 48 hr. The reaction mixture was then filtered to remove the unreacted copper and the filtrate added to water to precipitate the product. After recrystallizing from methanol, 6.6 grams (84%) of Compound 10 were obtained as light tan crystals: mp 170.0-171.5°C [lit. (8) mp 172°C]; ir (nujol) 1350, 1520 cm⁻¹ (NO₂); nmr (CDCl₃) & 7.7-7.9 (2H doublet, aromatic) and δ 8.2-8.4 (4H multiplet, aromatic).

Anal. Calcd C₁₀H₆N₂O₄: C, 55.05; H, 2.77; N, 12.85. Found: C, 55.00; H, 2.78; N, 12.88.

The above compound was also obtained in good yields when N-methylpyrrolidin-2-one or diethylformamide were substituted for DMAC.

3,3 - Bi 12H-benz (4,5) isoindolo (2,1-a) perimidin-12-one) (Compound 14). Phthalic anhydride (15 grams), powdered iron (3.4 grams), Compound 3 (4.3 grams) and acetic acid (50 ml) were refluxed for 4 hr and the reaction mixture filtered while hot. As the filtrate cooled, a red-brown precipitate formed, which was collected and recrystallized from acetic acid to yield 3.6 grams (67%) of Compound 7 as dark ruby crystals: mp 380-400°C dec; δ 7.6-7.8 (4H multiplet, aromatic), δ 7.6-7.8 (4H multiplet, aromatic), and & 8.3-8.5 (4H multiplet, aromatic).

Anal. Calcd C₃₆H₁₈N₄O₂: C, 80.29; H, 3.37; N, 10.40. Found: C, 80.15; H, 3.30; N, 10.45.

4.4'.5.5'-Tetraamino-1,1'-binaphthyl (Compound 15). Method A. Pulverized Compound 3 was mixed with palladium black (0.5 gram) in absolute alcohol (25 ml) and hydrazine hydrate was added dropwise over a period of 5 min, after which the reaction mixture was refluxed for 2 hr. After cooling, 250 ml of absolute ethanol was added and after the catalyst was removed by filtration, the volume was then reduced to 50 ml, the precipitate filtered and dried, yielding 0.5 gram (35%) of Compound 14 as a dark green amorphous solid: mp 200-230°C dec; ir (KBr) 3400, 3200 cm $^{-1}$, broad (NH₂); mass spectrum m/e 314 (molecular ion); nmr (DMSO-d₆) 53.7-4.1 (8H broad singlet, amine) and δ 8.3-8.7 (10H multiplet, aromatic).

Anal. Calcd C₂₀H₁₈N₄: C, 76.41; H, 5.77; N, 17.82. Found: C, 76.22; H, 6.14; N, 17.50.

Method B. A suspension of powdered Compound 3 in 40 ml of 0.5m sodium hydroxide solution and sodium bisulfite (3.0 grams) was refluxed for 15 min. After cooling, the mixture was filtered and the resulting black solid was recrystallized from nitrobenzene yielding 0.5 gram (35%) of Compound 15, identified by comparison of the ir and nmr spectra of the product of Method A.

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Synthesis of Tris(N,N-dimethyl-2-carbamoylethyl)phosphine Oxida

D. J. Daigle, ¹ T. L. Vigo, and L. H. Chance Southern Regional Research Laboratory, 2 New Orleans, La. 70119

Tris (N, N-dimethyl-2-carbamoylethyl) phosphine oxide was prepared by reacting phosphorus pentoxide with tris(2carboxyethyl) phosphine oxide in excess N, Ndimethylformamide. The compound was very hygroscopic and could only be recrystallized from a cold DMF-ethyl acetate-petroleum ether mixture.

Preparation of tris(N, N-dialkyl - 2 - carbamoylethyl)phosphine oxides have been reported in the patent literature but their physical properties, yields, and general characteristics of identification have not been described (5). These compounds were prepared by the following reaction (3):

$$\begin{array}{c}
O \\
\parallel \\
O \\
O \\
3(R.N-C-CH_2-CH_2)_3P=O + K_2HPO_3
\end{array}$$

The authors have prepared one of these compounds, tris (N, N-dimethyl-2-carbamoylethyl) phosphine oxide in quantitative yield by the Schindlbauer reaction, as shown by the following equation (6):

$$O = P(CH_1 - CH_2 - C_1 - OH)_3 + 1.5 P_2O_5 \xrightarrow{\text{reflux}} O$$

$$O = P(CH_1 - CH_2 - C_1 - CH_2)_3 + CO$$

(Because carbon monoxide is a side product, this reaction should be performed under a well-ventilated hood.)

The crude material isolated was hygroscopic. Because of this latter property and its solubility (soluble in polar solvents and insoluble in nonpolar solvents), the compound was difficult to purify, Initial purification involved dissolving the dark brown crystals in ethanol and adding

To whom correspondence should be addressed. One of the laboratories of the Sputhern Marketing and Nutrition Research Division, Agricultural Research Service, U.S. Department of Agripetroleum ether until a dark brown oil separated. The clear solution was decanted from the impure dark oil and evaporated to give light yellow crystals. Recrystallization was achieved by dissolving the yellow crystals in a minimum amount of N,N-dimethylformamide and adding a 2:1 by volume mixture of petroleum ether-ethyl acetate. The cloudy solution was refrigerated. The white needles were isolated the next day by filtering the solution under an argon atmosphere.

An ir spectrum of a chloroform solution of the compound showed the carbonyl absorption band at 5.93 μ (Vs) and the P=O absorption band at 8.6 μ (m). The proton nmr spectrum (Varian A-60A spectrometer) of a deuteriochloroform solution at 60 MHz showed a multiplet centered at δ 2.5 and two singlets, one at δ 3.03, the other at δ 3.13. The ratio of the multiplet to the methylgroups singlets was 3:2. The chemical shift between the methyl groups was due to their different environments in the planar structure (1, 2).

Experimental

Tris(N,N-dimethyl-2-carbamoylethyl)phosphine (1). Tris-(2-carboxyethyl)phosphine oxide (8.6 grams, 0.0323 mole) (4) was refluxed with phosphorus pentoxide (7.2 grams, 0.05 mole) in 125 ml of N.N-dimethylformamide for 15 hr under a well-ventilated hood. The resulting solution was cooled and the N,N-dimethylformamide (DMF-petroleum ether-ethyl acetate). Anal: Calcd for vacuum to give an oil which slowly crystallized. These dark crystals were dissolved in ethanol and enough petroleum ether added to separate a dark brown oil. The clear solution, after decantation and evaporation under vacuum, yielded 8.5 grams (76%). Mp 100-102°C (DMF-petroleum ether-ethyl acetate). Anal. Calcd for C₁₅H₃₀O₄N₃P; C, 51.85; H, 8.70; N, 12.09; P, 8.91. Found: C, 51.61; H, 8.81; N, 11.89; P, 8.84.

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